FOOD AND DRUG ADMINISTRATION

7500 Standish Place (HFV-210), Room N403 Rockville, MD 20855

VETERINARY ADVERSE DRUG REACTION, LACK OF EFFECTIVENESS, PRODUCT DEFECT REPORT

(Forward to address at left. Attach all correspondence that pertains to this reaction)

Form Approved: OMB No. 0910-0012 Expiration Date: December 31, 2001

Public reporting burden for this collection of information is estimated to average 1 hour per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to:

Food and Drug Administration 7500 Standish Place (HFV-210), Room N403 Rockville, MD 20855

An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control number.

NOTE: This report is required by law (21 CFR 510.300). Failure to report can result in withdrawal of approval of the application.							
1.	REPORT SOURCE AND ADDRESS (Mfr., Distr.)			2a. DATE REPORT RECEIVED	3a. TYPE OF REPORT		
					3-day Alert 15-day Alert		
				b. DATE SENT TO FDA	Periodic Report		
				NUMBER OF RAVO RETMEEN A AND I	3b. Initial Report		
				c. NUMBER OF DAYS BETWEEN 2a ANI	O b: Follow Up Report Of (Give Date)		
4.	NAME, ADDRESS AND PHONE NO. OF A (In confidence)	TTENDING VETERINARI	AN	5. NAME OR CASE IDENTIFICATION C (In confidence)	FOWNER		
	Name:						
	Street Address:						
	City: State:	ZIP:					
	Phone No. ()						
6.	TRADE NAME AND GENERIC NAME(S) O (Include dosage form and strength - Ex., tal	F ACTIVE INGREDIENT(b, 500 mg.)	S)	7a. NAME OF MANUFACTURER			
				b. NADA NO.			
8.	8. LOT NUMBER(S) 9. DOSAGE ADMINISTERED AND ROUTE			10. DATE(S) OF ADMINISTRATION			
(Ex. 250 mg., q 12 h, p.o.)							
11. ILLNESS/REASON FOR USE OF THIS DRUG			12. DRUG WAS ADMINISTERED BY				
The LEMESON FOR SOLE OF THIS BROSE			VETERINARIAN, STAFF OWNER, OTHER				
13.					SANIMALS		
a.	TREATED WITH DRUG b. REACTE	D c. DI	ED	a. SPECIES	b. BREED		
15.	CONCOMITANT MEDICAL PROBLEMS			c. AGE	d. WEIGHT		
				e. SEX			
				FEMALE MALE	PREGNANT NEUTERED		
				NY NEW ILLNESS DEVELOP OR DID INITI ECT DRUG STARTED?	AL DIAGNOSIS CHANGE AFTER		
GOOD FAIR POOR CRITICAL NO				YES (Explain)			
18. CONCOMITANT DRUG				IGS ADMINISTERED			
	NAME OF DRUG	ROUTE		DOSAGE REGIMEN	DATE(S) OF ADMINISTRATION		
FOR FDA USE ONLY							
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REACTION DATA DESCRIBE SUSPECTED ADVERSE REACTION: INCLUDE ALL SIGNS, RESULTS OF PERTINENT LAB TESTS, NECROPSY RESULTS, POSSIBLE CONTRIBUTING FACTORS, ETC. ALSO, INCLUDE IN THIS SECTION PRODUCT INEFFECTIVENESS AND PRODUCT DEFECTS SUCH AS CRACKED TABLETS, CLOUDY SOLUTION, ETC.				
a. ATTENDING VETERINARIAN'S LEVEL OF SUSPICION THAT DRUG CAUSED REACTION HIGH MEDIUM LOW NO ATTENDING VET. LENGTH OF TIME BETWEEN LAST ADMINISTRATION OF SUSPECT DRUG AND ONSET OF REACT	20b. WAS THERE EXTRA LABEL U NO YES (Exp. 22. DATE OF ONSET (Mo., day, yr.)	SE (ELU) INVOLVED? plain) 23. DURATION OF REACTION (Hrs., days, etc.)		
. WAS THE ADVERSE REACTION TREATED? NO YES (Describe treatment)	25. OUTCOME OF REACTION TO DED (Give date) REMAINS UNDER TREATM ALIVE WITH SEQUELAE RECOVERED UNKNOWN			
S. WHEN REACTION APPEARED, TREATMENT WITH SUSPECT RUG: HAD ALREADY BEEN COMPLETED DISCONTINUED DUE TO THE REACTION DISCONTINUED, REPLACE WITH ANOTHER DRUG DISCONTINUED, REINTRODUCED LATER CONTINUED AT ALTERED DOSE OTHER (Explain))		
B. DID ANIMAL(S) PREVIOUSLY REACT TO THIS DRUG?	NO YES UNKNOV NO YES UNKNOV NO YES UNKNOV	vn vn		
HAS THE ATTENDING VETERINARIAN SEEN SIMILAR REACTIONS TO THIS DRU NO YES (Describe treatment)	(If yes, give drug(s) and reaction if know	vn)		
	TURE OF INDIVIDUAL RESPONSIBLE	FOR ACCURACY OF REPORTED		

FORM FDA 1932 (2/99)